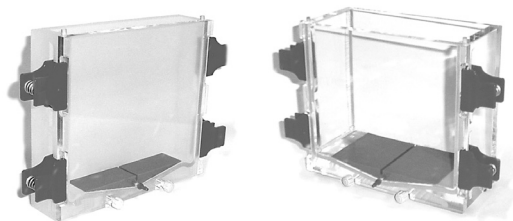


Hoefer SE615 and SE675

Multiple gel casters



Page finder

Safety warnings and precautions	viii
Introduction	1
Setting up the caster	5
Casting with divider plates	7
Pouring standard homogeneous gels	9
Pouring standard gradient gels	11
Pouring stacking gels	15
Removing polymerized gels	17
Troubleshooting	18
Care and maintenance	20
Appendix A: Gel identification numbers	21
Appendix B: References	21
Ordering information	22

Safety warnings and precautions

- Acrylamide is a neurotoxin.
- Wear gloves when handling acrylamide or polyacrylamide.
- Wear a dust mask when weighing acrylamide or preparing acrylamide solutions.

Protecting your equipment

To keep your instrument in excellent condition, please take the following important steps:

- Use water-saturated *n*-butanol for overlayering resolution gels during polymerization.
- Avoid using small metal spatulas to separate the plates. Spatulas may chip the edges of glass plates and prevent them from sealing.

After casting, you may need to use a wedge to separate one glass-and-gel sandwich from another. Slip a Wonder Wedge in between adjacent sandwiches to separate them.

Important! Most organic solvents including methanol, isobutanol, isopropanol, and even *n*-butanol will craze or cloud the acrylic plastic with prolonged exposure. Only use small amounts to overlay the gels.

Introduction

Gel sizes

The Hoefer® SE615 and SE675 Multiple Gel Casters are designed for casting vertical slab acrylamide gels of homogenous or gradient concentration.

Use the casters with 16 × 18 cm glass plates and 2 cm-wide spacers to cast 16 × 14 cm gels. Use 1-cm wide spacers to cast 16 × 16 cm gels for two-dimensional electrophoresis. The resulting gels are compatible with Hoefer SE400, SE600 and SE600 Chroma Vertical Slab Gel Electrophoresis Units.

Homogeneous and gradient gels

The SE615 and SE675 can be used for casting both homogeneous and gradient gels. Gradient gels are cast through the port at the bottom of the caster. When casting homogeneous gels, the results are most consistent when the gels are cast from the bottom. The V-shaped region at the bottom of the casting unit allows the gel solution to rise evenly, producing identical gels.

Standard casting

In the standard casting procedure, a gel is cast between two glass plates held apart by a pair of spacers. (See Fig 2 on page 5.) The result can be pictured as a “sandwich,” with the gel between two glass plates. When you set up a vertical slab gel electrophoresis unit, you place the glass and gel sandwich in the electrophoresis unit.

The maximum number of gels you can cast simultaneously using the standard casting procedure is shown in Table 1.

Table 1. Maximum number of standard gels cast

	0.75-mm spacers	1.0-mm spacers	1.5-mm spacers
SE615	11	10	10
SE675	4	4	4

Casting with divider plates

You may find it useful to double the number of gels run simultaneously in one electrophoresis unit. This can be accomplished by casting a “club sandwich.” A club sandwich contains a notched glass divider plate and a second pair of spacers between the standard plates, and thus forms two layers of gel between three glass plates (see Fig 3 on page 8.) Divider plates are available for the SE400 and SE600.

The maximum number of gels you can cast simultaneously, using divider plates (two gels per sandwich), is shown in Table 2.

Table 2. Maximum number of gels cast with divider plates

	0.75-mm spacers	1.0-mm spacers	1.5-mm spacers
SE615	14	14	12
SE675	6	6	4

See Fig 1 on page 4 to identify the components of the multiple gel casters. For information on parts, accessories and related equipment, see page 22.

Pouring second-dimension slab gels

When preparing second-dimension slab gels in the multiple casters, it is important to pour the slabs to the correct height.

If you are using IPG strips, you do not need a stacking gel. Pour the second-dimension gel to 0.5–1.0 cm below the top of the sandwich for standard casting, or 1 cm below the notch if you are casting with divider plates. This allows ample room to hold the IPG strips between the two glass plates. Use 1.0- or 1.5-mm spacers with IPG strips.

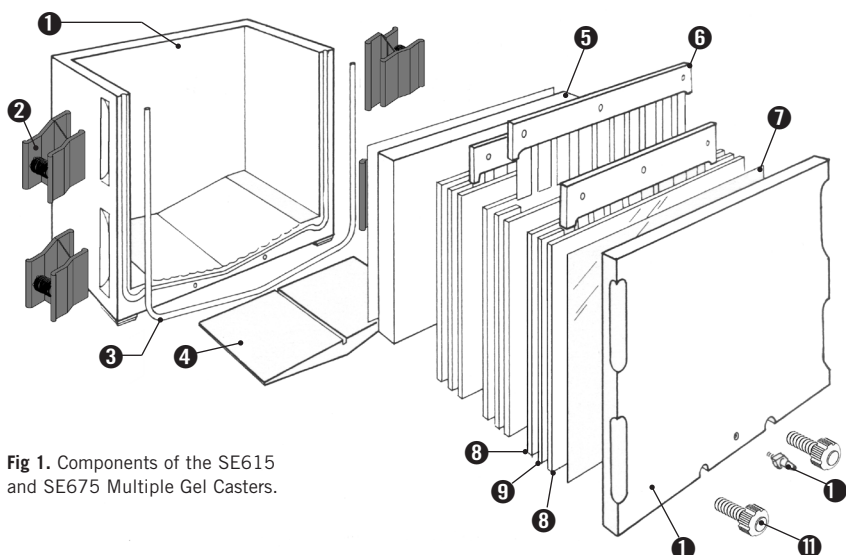


Fig 1. Components of the SE615 and SE675 Multiple Gel Casters.

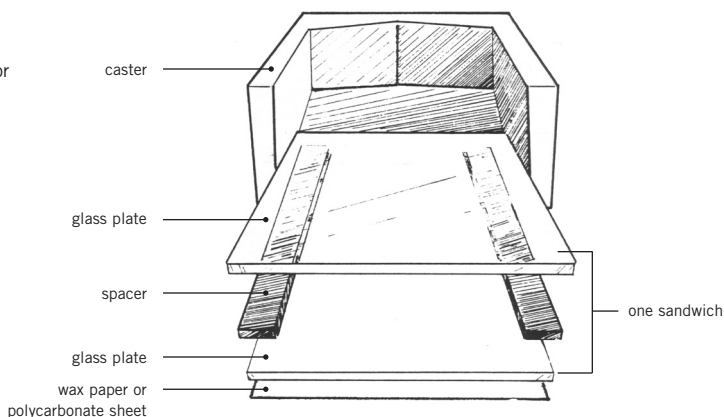
SE615 and SE675 components

no.	description
1	Caster body, includes faceplate
2	Spring clamp (4)
3	Foam cord gasket
4	Silicone displacement block*
5	Acrylic spacer block
6	Comb
7	Polycarbonate filler sheet
8	Glass plate
9	Spacer
10	Inlet port
11	Nylon thumb screw

*A small silicone displacement block is also available for the SE675.

Setting up the caster

Fig 2. Building a glass sandwich for standard casting.



Preparing the caster

1

If the faceplate is attached to the caster, remove it. Remove the four side clamps, loosen the two thumb screws, and slide off the faceplate.

2

Remove the foam cord gasket. Apply a light coat of Gel Seal lubricant and replace the gasket.

3

When casting gradient gels, attach the tubing from the gradient maker to the inlet port at the bottom of the faceplate. Homogeneous gels may also be cast through this port. If casting homogeneous gels from the top, insert the appropriate displacement block in the open, V-shaped region at the bottom of the chamber.

4

Place the open caster in a horizontal position with its back on the bench top.

Note: Do not block the red inlet port at the bottom of the faceplate.

Note: To make separation of polymerized gel sandwiches easier, place wax paper sheets in the caster between each set of glass sandwiches.

Note: When analyzing a large number of gels, you may find it useful to include gel identification numbers into the polymerized gel.

See the appendix on page 21 for directions on making gel identification tags. Lightly wet and place the tags in the lower corner of the gel sandwich during stack assembly.

Note: If you use polycarbonate sheets instead of wax paper between each sandwich, the caster holds fewer sandwiches.

Building the stack

1

Place a polycarbonate sheet in the caster so that 1/3 of the sheet extends out of the top.

You can use this sheet as a lever when inserting filler sheets after all sandwiches are in place. All remaining components fit flush against the bottom and sides.

2

Place a sheet of wax paper on the polycarbonate sheet. Build the first sandwich into the caster.

3

Place a sheet of wax paper on top of the sandwich and build the next sandwich.

4

Repeat step 3, alternating wax paper and gel sandwiches. Over the final sandwich, lay a polycarbonate sheet instead of wax paper so that the gel solution level is visible through the faceplate.

5

Place the faceplate on the caster and make sure the stack fits snugly, about 0.5 mm above the edge of the caster. If necessary, adjust the number of extra polycarbonate sheets, acrylic spacer blocks, glass plates and sandwiches to obtain a snug fit.

6

Make sure the spacers are straight and along the edges of the glass plates. Use the Spacer Mate to correct the alignment if necessary.

If necessary, take the faceplate off and make sure all edges of the stack components are flush. Press on those that are sticking out until all edges are flush and replace the faceplate on the caster.

7

Secure the faceplate with the four spring clamps and tighten the bottom thumb screws.

Casting with divider plates

To double the number of gels run simultaneously on the SE600 and related units, place a notched glass divider plate and another pair of spacers between the standard glass plates. The result is a “club sandwich,” with two layers of gel formed by the three glass plates.

To use divider plates, follow the directions in “Pouring standard homogeneous gels” on page 9 or “Pouring standard gradient gels” on page 11. When calculating solution volumes, use the formula on page 9. When stacking glass sandwiches, use the following procedure.

1

Stack glass sandwiches in the caster, using one club sandwich for each two gels you will cast. Fill extra space with acrylic blocks, glass plates and polycarbonate sheets.

2

Slide in one polycarbonate sheet against the back.

3

Slide in a glass club sandwich. (See Fig 3.)

To construct a club sandwich, slide in one rectangular glass plate and place spacers along each of the two side edges. Slide in a notched divider glass plate and place spacers along each of the side edges. Finally, slide in a second rectangular glass plate.

4

Place a polycarbonate sheet or wax paper on top of the club sandwich.

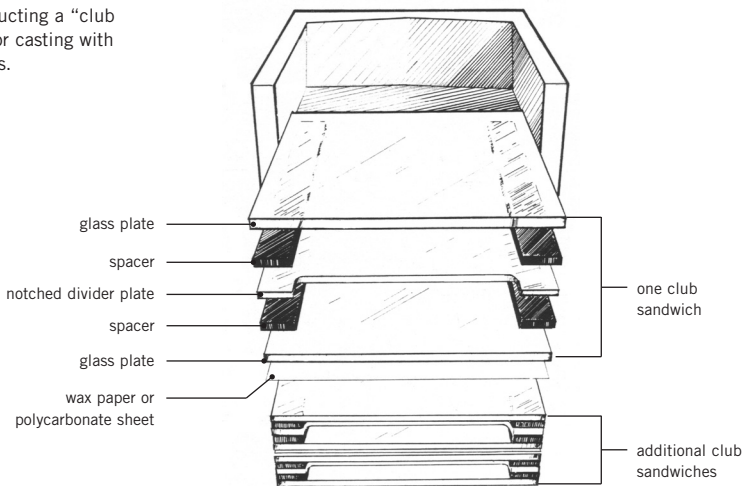
Repeat step 3, alternating wax paper and club sandwiches.

5

After the last club sandwich, add acrylic spacer blocks and extra polycarbonate sheets until the entire stack is approximately 0.5 mm above the rim of the box.

Note: For optimum visibility, place a polycarbonate sheet against the faceplate.

Fig 3. Constructing a “club sandwich” for casting with divider plates.



Warning! Acrylamide is a neurotoxin. Wear gloves when handling acrylamide or polyacrylamide. Wear a dust mask when weighing acrylamide or preparing acrylamide stocks.

Pouring standard homogeneous gels

Table 3. Gel casters and volume required

gel caster	required volume (ml)	gradient maker size*
SE615	190–500	SG500
SE675	60–150	SG100 or SG500

*Actual size depends on calculated volume.

1

Prepare and degas enough monomer solution to fill all sandwiches to the level of the notched plate.

The following formula allows extra solution to fill the space between the sandwich and the chamber:

Note: The 1.1 multiplier in this formula assures enough solution to fill dead volume.

$$\text{Monomer vol. (ml)} = \frac{\text{height (cm)} \times \text{width (cm)} \times \text{spacer thickness (cm)} \times \text{total number of gel sandwiches} \times 1.1}{1}$$

No stacking gel. Fill solution to just below the top of the notched plate. If air pockets form, remove with a pipette or syringe. Introduce a comb, at a slight angle, into each sandwich, taking care not to trap air under the teeth.

Stacking gel. Fill solution to 4 cm below the top of the rectangular glass plate. This height allows 1 cm of stacking gel below the wells. Pour the gel and apply an overlay. After the gel is set, prepare the stacking gel as described on page 15.

2-D electrophoresis. Fill solution to about 1.0 cm below the top of the rectangular glass plate. This height allows 4 to 5 mm of space for the IPG strip and an agarose seal. Overlay the separating gel as described in step 3.

Caution! Isobutanol crazes or clouds the acrylic caster walls, making it difficult to see the gels.

2

Add initiator and catalyst, and pour the solution into the glass sandwiches from the top.

The solution flows from one sandwich to another through the groove at the bottom of the caster and rises to the same level in all sandwiches. Pour slowly to avoid overfilling.

3

After reaching the final level, overlay each gel sandwich with 300 μ l of water-saturated *n*-butanol or buffer.

Use the same amount of overlay on each gel sandwich to assure that all the gels polymerize to the same heights. Pour the liquid gently, taking care not to disrupt the gel surface.

After polymerization is complete, remove the *n*-butanol and replace it with buffer or add stacking gel. Water-saturated *n*-butanol will cloud acrylic parts if used in large amounts or for long periods.

4

Allow the gels to polymerize for at least an hour. To remove the polymerized gels, see "Removing polymerized gels" on page 17. To add a stacking gel, see "Pouring stacking gels" on page 15.

Pouring standard gradient gels

See Fig 1 on page 4 for a diagram of the caster components. See “Setting up the caster” on page 5 for recommended solution volumes.

Setting up the gradient maker

1

Prepare a 50% glycerol displacement solution containing a small amount of bromophenol blue: approximately 75 ml for the SE615 and 35 ml for the SE675.

2

Calculate the amount of gradient solution you will need by referring to the formula on page 9.

For one-dimensional gels: Use a 12-cm separation gel with a 4-cm stacking gel.

For two-dimensional gels using IPG strips: Calculate the amount of solution needed to fill the sandwiches to approximately 0.5–1.0 cm below the top of the plates.

3

Set up the peristaltic pump. Using a graduated cylinder and water, adjust the flow rate so that the volume of the gradient separation solution plus the volume of the glycerol solution will be completely delivered in 10–15 minutes.

4

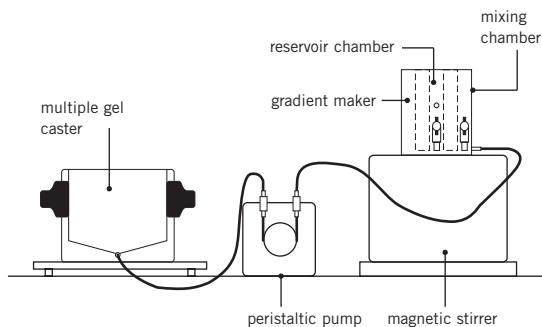
Choose a gradient maker that holds no more than four times the total volume of gradient solution to be poured. Check that the gradient maker is clean and that the outlet and mixing port are free of polymerized acrylamide.

5

Assemble the gradient making system. (See Fig 5.)

- Close all gradient maker valves and place a stir bar in the mixing chamber, the one with the tube connector port.
- Attach one end of a piece of tubing to the outlet of the gradient maker.
- Run the other end of the tubing through the peristaltic pump and attach it to a tubing connector.
- Use a second piece of tubing to attach the tubing connector to the red inlet port at the bottom of the caster.

Fig 5. Gradient making system connections.



Warning! Acrylamide is a neurotoxin. Wear gloves when handling acrylamide or polyacrylamide. Wear a dust mask when weighing acrylamide or preparing acrylamide stocks.

Pouring gradient separation gels

1

Prepare and degas the two monomer solutions for the gradient maker.

Add glycerol or sucrose to the solution of higher acrylamide concentration to stabilize the gradient as the solution is pumped in. Decrease the concentration of initiator so that polymerization occurs from top to bottom. This minimizes convective mixing due to the heat generated by polymerization.

2

Close both the mixing port and outlet port, if appropriate, on the gradient maker. Clamp the tubing from the outlet when using the SG500.

3

Add initiator and catalyst, and immediately pour the light (low concentration) acrylamide solution into the mixing chamber—the chamber with the outlet port. Open the mixing valve slightly to allow the connecting channel to fill and force out air bubbles.

Close the valve again and pour the heavy (high concentration) acrylamide solution into the reservoir chamber.

4

Start the magnetic stirrer and unclamp or open the outlet valve. Start the pump and open the mixing valve.

5

When *almost all* the acrylamide solution is drained from the gradient maker, stop the pump and close the mixing valve. Tilt the gradient maker towards the outlet side and remove the last few milliliters of mix. *Do not allow any air bubbles to enter the tubing.*

6

Add the glycerol displacement solution to the mixing chamber and start the pump. Make sure no bubbles are introduced. Pump until the bottom of the caster is filled with displacement solution to just below the glass plates, then turn off the pump. Clamp off the tubing to the red inlet port of the caster.

7

Overlay each separate sandwich with 300 μ l of water-saturated *n*-butanol or buffer.

Use the same amount of overlay on each gel to assure that all the gels polymerize to the same height. Carefully and gently pipette the liquid, taking care not to disrupt the gel surface.

After polymerization, remove the *n*-butanol. Replace it with buffer or add a stacking gel.

8

Allow the gels to polymerize for at least an hour.

To remove the polymerized gels, see “Removing polymerized gels” on page 17. To add a stacking gel, see “Pouring stacking gels” on page 15.

Note: Isobutanol clouds the acrylic, making it difficult to see the gels.

Pouring stacking gels

You may add stacking gels to the entire set of polymerized gels—homogeneous or gradient—while the sandwiches are still in the casting box, or you may add stacking gel to individual gels immediately prior to use.

For optimal resolution, add the stacking gel at the last minute to prevent diffusion of buffers between the two gel layers.

1

If you are pouring stacking gels later, skip to step 1 under “Removing polymerized gels” on page 17.

If you are pouring stacking gels now, calculate the volume of stacking gel monomer solution needed by referring to the following formula:

$$\text{Monomer vol. (ml)} = \frac{\text{height (cm)}}{\text{cm}} \times \frac{\text{width (cm)}}{\text{cm}} \times \frac{\text{thickness (cm)}}{\text{cm}} \times \frac{\text{total number of stacking gels}}{\text{cm}}$$

Where:

- Height is the distance between the separating gel and the top of the plates.
- Width is the distance between the spacers.

2

Rinse off the water-saturated *n*-butanol overlay with distilled water or Tris buffer and invert the caster to drain it. Repeat this 2–3 times.

3

Degas the stacking gel monomer solution and add catalyst and initiator.

Note: Oxygen inhibits gel polymerization. Do not trap air bubbles underneath the comb teeth.

Note: Use gels with stacking gels immediately. Do not store.

4

Replace the caster in a vertical position. Make sure the surface of the separating gel is free of liquid. Use a pipette to fill each sandwich individually with stacking-gel monomer solution.

5

Hold a comb, attached to a comb back, at a 45° angle and centered within the sandwich. Insert one end into the gel so that the end tooth is almost completely inserted. Slowly lower the remaining comb teeth in one by one, rotating the comb gradually downward until it is in a horizontal position.

Refer to Fig 1 on page 4 for the final position of the comb. Repeat this for each sandwich.

6

Allow the stacking gels to polymerize for at least one hour. To remove the polymerized gels, see “Removing polymerized gels” on page 17.

Note: The SDS buffer makes the glass plates slippery.

Note: Do not use metal spatulas to separate the sandwiches. Narrow metal spatulas often chip the edges of glass plates, making the plate ineffective for sealing into the electrophoresis tank.

Note: Separate the sandwiches from one another before placing the gels in the refrigerator.

Removing polymerized gels

1

After the gels have polymerized, place the caster in a horizontal position. If you poured stacking gels, leave the combs in place. Remove the faceplate.

2

Tip the caster to pour off the glycerol solution.

3

Slide the stack of sandwiches, acrylic blocks, and polycarbonate sheets out of the caster. Remove the acrylic blocks and polycarbonate sheets from the top of the stack.

4

Carefully remove individual glass and gel sandwiches from the stack. If necessary, insert a Wonder Wedge between adjacent sandwiches to separate them.

Work slowly and cautiously when prying apart adjacent sandwiches after polymerization.

5

Rinse the caster and polycarbonate sheets, as described in “Care and maintenance” on page 20.

To store gels for future use

Rinse the individual sandwiches with distilled water. Fill cassettes with gel storage buffer. Wrap gel sandwiches you are not using at this time in plastic wrap and store them in the refrigerator.

To use gels immediately

1

Gently remove the combs by pulling them upward and out of the gel sandwich.

2

Fill the wells with SDS Electrophoresis buffer.

3

Load samples and complete assembly. Refer to the user manuals that come with SE600 or SE400 Series electrophoresis systems.

Troubleshooting

symptom	possible cause	recommended action
1. Gels adhere to glass plates when opening sandwich.	Dirt, grease or fingerprints on plates.	Soak plates in a strong laboratory detergent; rinse well in distilled water. Handle with gloves only.
2. Gels cast simultaneously are different sizes.	Different amounts of overlay were used on the separation gels before polymerization.	Use the same amount of overlay on all separation gels. Add the overlay as rapidly as possible.
3. Caster leaks.	Gasket leaks.	Apply a light film of Gel Seal to the gasket each time the unit is used.
	Gasket damaged.	Check the foam gasket for nicks or wear and replace if necessary.
	Stack too tall.	Remove filler plates or gel sandwiches until the stack top is just below the level of the caster wall.
4. Gel heights uneven.	Bubbles trapped under gel sandwiches.	Pour the monomer solution into one sandwich and allow the groove in the plug to evenly distribute the solution.
	Polymerized gel in groove.	Make sure the groove in the triangular plug is clean and clear of material.
	Insufficient time for gel levels to stabilize.	Wait one minute before overlaying the gels. Add the same amount of overlay to each sandwich.
5. Sample wells damaged or leak.	Air bubbles under comb teeth.	Remove air pockets before inserting combs. Slide comb into solution at an angle. If comb must be removed, add more monomer solution before reinserting the comb.
	Insufficient polymerization.	Allow the gel to set for a minimum of 1 hour.
	Comb removed too abruptly.	Slowly remove the comb at a slight angle to prevent damaging the gel.
	Poor polymerization.	Degas stacking gel solution. Increase catalysts up to 0.1% v/v TEMED, 0.1% v/v APS.
6. Gel sandwiches difficult to separate.	No wax paper between glass plates.	Place wax paper between sandwiches in the stack.
7. Uneven gradient gels.	Uneven layering.	Add sucrose or glycerol to the heavy monomer solution. Add a small amount of bromophenol blue to the heavy solution to track gradient formation. Decrease the pump rate.

symptom	possible cause	recommended action
8. No polymerization of SDS gel (or incomplete polymerization.)	Insufficient APS or TEMED.	Increase both APS and TEMED by 30–50% APS solution is old. Make up fresh APS each day.
	APS stock is wet.	APS is hygroscopic. Open a fresh bottle.
	O ₂ in gel solution.	Degas at least 10 minutes.
	Solutions at low temperature.	Make sure all solutions are at room temperature (20–30 °C).
	TEMED is old.	Use new TEMED.
9. Gel too soft.	Not enough crosslinker.	Crosslinker should be 2.6% C for standard SDS gels where $\%C = \frac{\text{g bis}}{(\text{g monomer} + \text{g bis})} \times 100$
10. Gel is brittle.	Too much bis.	See #9 above.
11. Gel is white.	Too much bis.	Check concentrations of solutions. See #9 above.
12. Gel contains swirls, polymerization artifacts.	<i>Too much catalyst:</i> gel polymerized in < 10 min.	Reduce both APS and TEMED by 25%.
	<i>Not enough catalyst:</i> gel polymerized in > 50 min.	Increase both APS and TEMED by 50%. Also see #8.
	Solutions not mixed.	Mix thoroughly after adding TEMED.
13. Bands are diffuse or broad.	Sample doesn't contain same buffer as stacking gel.	Use the same buffer for the sample as for the stacking gel.
	Too much TEMED or APS.	Reduce concentrations by 25%.
	SDS or sample buffer is old.	Use fresh solutions.
	Poor interface between separation gel and stacking gel.	Remove all liquid from the surface of the separation gel before adding the stacking gel solutions.
	Incomplete polymerization.	See #8.
14. Protein mobilities not consistent.	Aged gels or acrylamide.	Do not store liquid acrylamide more than 3 months. Use gels within 1–2 weeks of casting. Use gels with stackers immediately.
	Gas in gel.	Degas gel solutions at least 10 minutes.
15. Heavy background during silver staining.	Acrylamide or bis contain acrylic acid.	Use reagents specified as electrophoresis purity.
	Water is impure.	Use only double-distilled water.

Care and maintenance

- Do not autoclave or heat any part above 45 °C.
- Do not expose the caster or its parts to organic solvents.

Cleaning

Rinse the caster, faceplate, silicone plugs and polycarbonate sheets in dilute detergent and rinse with distilled water. Allow the unit to air dry completely.

Appendix A: Gel identification numbers

For positive identification of gels, label each slab by incorporating a small label printed on thin filter paper in the bottom corner of the gel. Use a carbon typewriter ribbon, photocopier or laser printer to make these labels, since many liquid-based inks are electrophoresed off paper during an SDS electrophoresis run.

A variety of numbering schemes are possible. In our experience the easiest uses three parts as follows:

- An upper-case letter to identify the investigator or an extended gel series.
- A two- or three-digit serial number to identify the slab gel batch.
- A lower-case letter to identify a gel in the batch. Since a maximum of 14 gels can be made in a batch, use the letters a–n.

The resulting numbers, in the format A63a, A63b..., etc., provide a useful system for keeping track of and cross-indexing experiments and gel production.

Appendix B: References

Laemmli, U.K. 1979. *Nature* (London) **227**: 680–685.

SDS-PAGE and IEF Handbook.

Ordering information

product	quantity	code number
SE615 Multiple Gel Caster Kit, 10 gels Includes 20 glass plates, space-saver plate, 5 filler sheets, 100 sheets of wax paper, and Spacer-Mate alignment template. (Order combs and spacers separately.)	1	SE615
SE675 Multiple Gel Caster Kit, 4 gels Includes 8 glass plates, 3 space-saver plates, 5 filler sheets, 100 sheets of wax paper, and Spacer-Mate alignment template and filler plugs. (Order combs and spacers separately.)	1	SE675
Foam cord gasket, 61 cm × 4.5 mm OD	1	SE208
Red inlet port	4	XP010
Red spring clamps	4	SE253
Glass plates, 18 × 16 cm	2	SE6102
Notched divider glass plates, 18 × 16 cm	1	SE6102D
Acrylic block, 11 mm thick, 18 × 16 cm	1	SE612
Polycarbonate filler sheets	5	SE613
Wax paper, precut sheets, 18 × 16 cm	100	SE614
Spacer-Mate template for aligning spacers	3	SE6119SM
Nylon thumb screws	12	SE6003U-2
Silicone filler plug, SE615	1	SE618
Silicone filler plug set, SE675. Includes 1 large and 1 small plug.	1	SE678
Gel Seal, ¼ oz. tube	1	SE6070

Related products

SE100 PlateMate, plate washer and storage unit	1	SE100
SG100 Gradient Maker, 100 ml total volume	1	SG100
SG500 Gradient maker, 500 ml total volume	1	SG500
Wonder Wedge	1	SE1514

Spacers

length (cm)	thickness (mm)	width (cm)		
16	0.75	2	2	SE6119-2-.75
16	1.0	2	2	SE6119-2-1.0
16	1.5	2	2	SE6119-2-1.5
16	1.0	1	2	SE6118-2-1.0
16	1.5	1	2	SE6118-2-1.5

Combs

number of wells	thickness (mm)	width (mm)	quantity	code number
10	0.75	8.3	1	SE511-10-.75
10	1.00	8.3	1	SE511-10-1.0
10	1.50	8.3	1	SE511-10-1.5
12	0.75	7.6	1	SE511-12-.75
12	1.00	7.6	1	SE511-12-1.0
12	1.50	7.6	1	SE511-12-1.5
15	0.75	5.7	1	SE511-15-.75
15	1.00	5.7	1	SE511-15-1.0
15	1.50	5.7	1	SE511-15-1.5
20	0.75	4.1	1	SE511-20-.75
20	1.00	4.1	1	SE511-20-1.0
20	1.50	4.1	1	SE511-20-1.5
28 ^a	0.75	2.7	1	SE511-28-.75
28 ^a	1.00	2.7	1	SE511-28-1.0
28 ^a	1.50	2.7	1	SE511-28-1.5

^aComb depth 15 mm; all others 25 mm.

Preparative combs

These combs are 25 mm deep, adjustable to 10 or 15 mm.

no. of wells prep/ref	thickness (mm)	width (mm) prep/ref	quantity	code number
1/1	0.75	121/6	1	SE511-R-.75
1/1	1.00	121/6	1	SE511-R-1.0
1/1	1.50	121/6	1	SE511-R-1.5
1/2	0.75	113/6	1	SE511-DR-.75
1/2	1.00	113/6	1	SE511-DR-1.0
1/2	1.50	113/6	1	SE511-DR-1.5

Adjustable comb back

1

SE511-BKA

Required to convert any 25-mm deep comb to 10 or 15 mm depth.

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